Run on:	OM nucleic	
March 7, 2007, 12:57:57; Search time 567 Seconds	OM nucleic - nucleic search, using sw model	AND THE TOTAL TOTA

653.489 Million cell updates/sec

Scoring table: IDENTITY_NUC Gapop 10.0 ,	Sequence:	Title: US.
IDENTITY NUC Gapop 10.0 , Gapext 1.0.	1 tgccatgtgatactccaggaactatttccaatgatgatga 50	US-10-751-113-3

Total number of hits satisfying chosen parameters:

5620219 seqs, 3705283702 residues

Searched:

11240438

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries Minimum DB seq length: 0
Maximum DB seq length: 2000000000 .

Database : N_Geneseq_200701:*
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2: geneseqn2000s:*
3: geneseqn2001as:*
5: geneseqn2001bs:*
6: geneseqn2002as:*
7: geneseqn2002as:*
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geneseqn2001bs:*
geneseqn2001bs:*
geneseqn2002bs:*
geneseqn2002bs:*
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geneseqn2003ds:*
geneseqn2004as:*
geneseqn2004bs:*
geneseqn2005s:*
geneseqn2005s:*

00 0000

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

1	Result
50	Score
50 100.0	% Query Match
50 8	ult Query No. Score Match Length DB ID
50 8 ABX11900	
Abx11900 Human AIB	Description

45	.44	43	42	41	40	39	88	37	36	35	34	ü	32	31	30	29	28	27	26	25	24	23	22	21	20	19	18	17	16	15	14	13	12	11	10	9	æ	7	0	Ŋ	4	ω	2
23.6	23.8	23.8	23.8	23.8	23.8	24.2	24.2	24.2	24.4	24.4	24.4	24.4	24.8	25	26.2	26.2	26.4	26.4	26.4	26.4	27.6	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8	27.8
47.2	47.6	47.6	47.6	47.6	47.6	48.4	48.4	48.4	48.8	48.8	48.8	48.8	49.6	50.0	52.4	52.4	52.8	52.8	52.8	52.8	55.2	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6
11614	187851	169659	44528	10199	864	110000	110000	1664	171936	171936	3361	1179	888	35	4860	4621	268685	264965	86453	403	34	12414	8207	8207	7994	7935	7923	7116	7088	6845	6835	6835	6835	6832	6754	6754	6754	6754	6754	4789	4495	4263	4239
14	14	12	6	6	œ	14	14	4	12	6	2	4	13	8	N	N	6	12	14	9	æ	14	10	10	14	15	15	10	15	10	14	14	N	0	14	14	14	10	4	N	14	12	14
AEA61172	ADZ13735	ADQ59434	ABA03041_29	ABQ71072	ACA36808	AEB42737_12	AEB39174_07	AAF74192	ADN16205	ABS56565	AAX80993	AAK87368	ADR 634 01	ABX11898	AAX26000	AAV99919	ABS56563	ADN16203	ADY25730	ACH16713	ABX11899	·AEA47654	ADF81690	ADF81691	ADY16115	AEH88176	AEH88178	ADE53851	AEF22148	ADE76189	AED18120	ADY19927	AAV99915	AAD30439	ADY16117	ADY16113	ADY15855	ADK67037	AAF26490	AAX80992 .	AEC82913	ADJ92834	ADV43475
Aea61172 Human HRA	Adz13735 Human can	Adq59434 Human can	Continuation (30 o	Abq71072 Listeria	Aca36808 Prokaryot	Continuation (13 o	Continuation (8 of	Aaf74192 DNA encod				Aak87368 Human imm	-	Abx11898 Human AIB			Abs56563 Human SUL		Ady25730 SULF rela	Ach16713 Human adu	т:	Aea47654 Nucleotid	_	•			Human					DNA			DNA			7 Gene		Human e		Human	Adv43475 Human psy

ALIGNMENTS

RESULT 1
ABX11900
ID ABX1
XX
AC ABX1
XX
DT 16-M
XX 16-MAY-2003 (first entry) ABX11900; ABX11900 standard; cDNA; 50 BP.

Human AIB1 delta3 isofrom mRNA exon 2/exon 4 junction

ovarian cancer; pancreatic siRNA; small inhibitory RNA; cytostatic; intestinal cancer; head and neck cancer; metastatic tissue cancer; neuroblastoma; Human; ss; exon 2/exon 4 junction; amplified in breast cancer; chromosome 20q; transcription factor; antisense gene therapy; transgenic; breast cancer; AIB1 delta3 isoform. cancer; prostate cancer; stomach cancer; amplified in breast

Homo sapiens.

WO2003003904-A2

16-JAN-2003

03-JUL-2002; 2002WO-US021066.

05-JUL-2001; 2001US-0302648P

(GEOU) UNIV GEORGETOWN MEDICAL CENT.

Reiter R, Wellstein A;

WPI; 2003-221517/21

New isolated AIB1 isoform as co-activator that potentiates transcriptional activity of nuclear hormone receptors, useful for the diagnosis, prevention and/or treatment of cancers, in particular breast

Example 6; Fig 9; 47pp; English

neuroblastoma, ovaries, pancreas, prostate, stomach, and in particular cancer of the breast. The gene for AIB1 is located on chromosome 20q. The present sequence is the junction region of the AIB1 deltral isoform mRNA for exons 2 and 4 (unique to the deltal isoform), which was targeted by detecting binding), a pharmaceutical composition (comprising an agent that specifically binds to the AIB1 isoform, and prevents a co-activation of a transcription factor when administered to a patient), an SIRNA nucleic acid), an anti-AIB1 isoform antibody (or antibody fragment), detection of cancer in a patient (comprising contacting a biological sample obtained from the patient with at least one chemical substance that specifically binds to the AIB1 isoform or the nucleic acid, and isolated nucleic sequences that encodes the isoform, a vector that contains the nucleic acid, a recombinant cell that comparising chemical acid, a diagnostic kit for the detection of cancer (comprising chemical substances that are specifically reactive to the AIB1 isoform or the treatment of disorders associated with the co-activator isoforms, such as cancers of the intestines, head and neck, metastatic tissue, compositions comprising the AIB1 isoform or nucleic acids that bind to the AIB1 isoform, are useful for the diagnosis, prevention and/or activator protein, a transgenic animal comprising a recombinant gene that encodes the AIB1 isoform and creating the transgenic animal. The The invention relates to an isolated isoform of an amplified in breast cancer 1 (AIB1) transcriptional co-activator. Also included are an (small inhibitory RNA) that inhibits expression of a transcriptional co-The

NO 1829 are provided.

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RESULT 2
ADV43475
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Best Local S
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                                                                                                                                                           The invention relates to a new microarray which comprises probes for genes involved in psychoneuroendocrinimmune (PNI) activity. The microarray is useful in diagnosing a condition associated with PNI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               04-JUN-2004; 2004WO-US017686
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Homo sapiens.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 microarray; psychoneuroendocrinimmune; chronic fatigue; non-insulin dependent diabetes; allergy; immune disorder; cancer; neoplasm; infection; expressed sequence tag; ss.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human psychoneuroendocrinimmune expressed sequence tag SEQ ID NO 1103
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10-MAR-2005 (first entry)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADV43475;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Sequence 50 BP; 18 A; 9 C; 11 G;
                                                                                                                                                                                                                                                                                                                                                                                                 New microarray comprising probes for genes involved in psychoneuroendocrinimmune (PNI) activity, useful in diagnosing a condition associated with PNI activity, e.g., inflammatory or information of the property of the condition associated with PNI activity, e.g., inflammatory or information of the property of the prop
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  HPI;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Nicholson A,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            04-JUN-2003; 2003US-0475915P
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  W02004108899-A2
                            psychoneuroendocrinimmune gene expressed sequence tag. Note the specificatio mentions SEQ ID NO of up to 3314 but only sequences up to
                                                                                                                                                                                                                                                                                                     Claim 1; SEQ ID NO 1103; 254pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (USSH ) US DEPT HEALTH & HUMAN SERVICES.
                                                                                               cancer and infection. The present sequence represents a
                                                                                                                               activity, such as CFS, type-2 diabetes, allergic condition, inflammation
                                                                                                                                                                                                                                                                                                                                                                            diseases
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  2005-031682/03
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           1 TGCCATGTGATACTCCAGGACAAGGGAAAAACTATTTCCAATGATGATGA
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Vernon
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                                                                                                                                                                                                                                                                                                                                                                                                               or infectious
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Title:	Run on:	OM nucleic -
US-10-751-113-3	March 7, 2007, 13:22:36 ; Search time 2477 Seconds (without alignments) 1395.242 Million cell updates/sec	OM nucleic - nucleic search, using sw model

Scoring table: IDENTITY_NUC
Gapop 10.0 , Gapext 1.0

Perfect score: 50
Sequence: 1 tgccatgtgatactccagga.....actatttccaatgatgatga 50

Total number of hits satisfying chosen parameters: 15137082

7568541 seqs, 34560148153 residues

Searched:

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

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7: gb_sts:*
8: gb_sv:*
9: gb_un:*
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13: gb_htg:*
14: gb_om:*
15: gb_ba:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

2	Result
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55.6	% Query Match
480 2 4495 2	Query Score Match Length DB ID
N N	8
CQ712958 Sequence CS162240 Sequence	Description

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7.6 55	7.6 55	27.6 55	σ	27.6 55	σ	6	7.6 55	6	.6	7.6 55	.0	σ	-	œ	80		.8	80	80	œ	œ	89		œ	.8	œ		80					20	œ	œ	œ	œ	œ	7.8 55	7.8 55	7.8 55	/.8 55
.2 110000	.2 110000	5.2 110000	.2 110000	.2 110000	.2 110000	.2 110000	. 2 110000	.2 110000	•	.2 110000	.2 90750	'n	.2 8	.6	6	.6	6	.6 7433	.6 7433	55.6 6835	•			.6 6835	σ	δ	6	თ	6	σ,	6	י יס	ν.	σ.	σ	0	6	0	σ	.6 4807	.6 4668	.6 449
•	4	4	4	4	4	4	4	4	4	4	4	4	4	N	2	2	N	Ŋ	ហ	u	ν	N	N	N	Ν	N	u	2	N	2	2	ν ι	v	N	N	N	N	N	ហ	տ	v	U
AP008218_213	AP008218_195	AP008218_067	AP008216_091.	AP008215_152	AP008213_099	AP008211_029	P008210_147	AP008210_134	P008209_210	AP008209_209	AP003797	CNS07YPG	AP003290	AX780090	AX780089	CS041367	CS032415	BC122547	BC119001	AF012108	CS045179	CS036227	AR316837	BD192354	AX384818	CQ730100	AF036892	AR908661	DD219303	CS050236	CS041369	CS041365	CS041107	CS032417	CS032413	CS032155	BD313717	AX821998	AB209863	BC092516	AF016031	AFULU221
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ion (214	ion (196	ion (68 o	ion (92 o	ion (153	ion (100	ion (30 o	ion (148	ion (135	ion (211	ion (210	Oryza sat	Oryza sat	Oryza sat	Sequence	Sequence	Sequence	Sequence		Homo sapi	Homo sapi	Sequence	Sequence	Sequence	AIB1, a s	Sequence	Sequence	Homo sapi	Sequence	GENE EXPR	Sequence	Sequence	Sequence	Segrence	Sequence	Sequence	Sequence	Antisense		Homo sapi	Ното варі	Homo sapi	TORO PAPT

ALIGNMENTS

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ORGANISM	KEYWORDS	VERSION	ACCESSION	DEFINITION	LOCUS	CQ712958	RESULT 1
Homo sapiens (numan)		CQ712958.1 GI:42273815	CQ712958	Sequence 57884 from Patent WO02070737.	CQ712958 480 bp DNA linear PAT 03-FEB-2004		

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Primates; Haplorrhini; Catarrhini; Hominidae; Homo.

REFERENCE 1

AUTHORS Liew,C.C., Marshall,W.E. and Zhang,H.
TITLE Compositions and methods relating to osteoarthritis
JOURNAL Patent: WO 02070737-A 57884 12-SEP-2002;
Chondrogene Inc. (CA)
FEATURES Location/Qualifiers
source /organism="Homo sapiens"
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Query Match 55.6%; Score 27.8; DB 2; Length 480; Best Local Similarity 82.1%; Pred. No. 16; Matches 32; Conservative 0; Mismatches 7; Indels 0; Gaps 0;

CS162240
LOCUS
CS162240
LOCUS
CS162240
DEFINITION
Sequence 481 from Patent WO2005083429.
ACCESSION
CS162240
VERSION
CS162240
VERSION
CS162240.1 GI:76060367
KEYWOEDS
SOURCE
ORGANISM Homo sapiens (human)
ORGANISM Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Euarchontoglires; Primates; Haplorrhini;
Catarrhini; Hominidae; Homo.

REFERENCE 1
AUTHORS Wang, Y.
TITLE Breast cancer prognostics
JOURNAL Patent: WO 2005083429-A 481 09-SEP-2005;
FEATURES Location/Qualifiers
SOURCE 1..4495

/organisms Homo sapiens // organisms Homo sapiens // mol_type="unassigned DNA" //db_xref="taxon:9606"

ORIGIN

Query Match 55.6%; Score 27.8; DB 2; Length 4495; Best Local Similarity 82.1%; Pred. No. 15; Matches 32; Conservative 0; Mismatches 7; Indels 0; Gaps 0;

REFERENCE AUTHORS TITLE REFERENCE AUTHORS SOURCE ORGANISM RESULT 3 AF010227 KEYWORDS VERSION DEFINITION POCUS FEATURES ACCESSION TITLE JOURNAL JOURNAL PUBMED gene source Sa complete cds. AF010227 Submitted (25-JUN-1997) Pharmacology and Molecular Toxicology, University of Massachusetts Medical School, 55 Lake Avenue North, Worcester, MA 01655-0126, USA 2 (bases 1 to 4495) Chen, J.D., Li, H. and Gomes, P.J. RAC3, a steroid/nuclear receptor-associated coactivator that is related to SRC-1 and TIF2 $\,$ Homo sapiens Homo sapiens receptor-associated coactivator 3 (RAC3) mRNA, Direct Submission 9238002 Proc. Natl. Acad. Sci. U.S.A. 94 (16), 8479-8484 (1997) Li,H., Gomes, P.J. and Chen, J.D. Mammalia; Eutheria; Euarchontoglires; Primates; Haplorrhini; Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Homo sapiens (human) AF010227.1 GI:2318005 AF010227 Catarrhini; Hominidae; Homo. (bases 1 to 4495) /translation="msglgenldplasdsrkrklpcdtpgggltcsgekrrredgesky ieelaelisanlsdidhokdrdkeskaliveyerdirqikgedlvatsyval teelaelisanlsdidhokdrdkeskaliveyerdiqikgedlvatsyval teeldrkdflkgilpkstvngvswtnetorokkshtencrmlmktphdiledinaspems /product="receptor-associated coactivator /protein_id="AAC51663.1" /db_xref="GI:2318006" associated coactivator 3; similar to the mouse p300/CBP/co-integrator protein (p/CIP) and the activator which includes steroid receptor coactivator 1 (SRC-1), transcriptional intermediate factor 2 (TIF2), and receptor >histone acetyltransferase activity; member of the steroid/nuclear receptor-associated coactivator family Location/Qualifiers PSSYQNNNYGLNMSSPPHGSPGLAPNQQNIMISPRNRGSPKIASHQFSPVAGVHSPMA SSGNTGNHSFSSSSLSALQAISEGVGTSLLSTLSSPGPKLDNSPNMNITQPSKVSNQE QRYETMQCFALSQPRAMMEEGEDLQSCMICVARRITTGERTFPSNPESFITRHDLSGK VVNIDTNSLRSSMRPGFEDIIRCIQRFFSLNDCQSWSQKRHYQEAYLNGHAETPVYR of retinoid receptors (ACTR) " GCNSSVGGMSMSPNQGLQMPSSRAYGLADPSTTGQMSGARYGGSSNIASLTPGPGMQS 1. .4495 /db_xref="taxon:9606" /cell_line="HeLa" organism="Homo sapiens" fsladgtivtaqtksklfrnpvtndrhgfvsthflqreqngyrpnpnpvgqgirppma /codon_start=1 note="transcriptional coactivator with intrinsic" gene="RAC3" gene="RAC3" mol_type="mRNA" 4495 bp IRNA linear PRI 13-AUG-1997

Run on: March 7, 2007, 13:32:22; Search time 5206 Seconds
(without alignments)
595.861 Million cell updates/sec OM nucleic - nucleic search, using sw model

US-10-751-113-3 50

Title:
Perfect score:
Sequence: 1 tgccatgtgatactccagga.....actatttccaatgatgatga 50

Scoring table: IDENTITY_NUC Gapop 10.0 , Gapext 1.0

Searched:

Total number of hits satisfying chosen parameters: 107170430

53585215 seqs, 31020513797 residues

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database : EST:

gb_est1:*
gb_est3:*
gb_est4:* gb_est5:*

gb_est6:*
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gb_est11:*
gb_est13:*
gb_est13:*
gb_est3:*
gb_gss1:*
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gb_gss3:*
gb_gss3:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result * Query

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	51.6	51.6	52.0	52.4	52.4	52.4	52.4	N	52.4	52.4	52.4	52.4	52.4	52.4	52.4	52.4	52.8	53.2	53.6	55.2	55.2	55.2	55.2	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	55.6	S	S	S	55.6	55.6	55.6	55.6	55.6	56.4	Match Length
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	BF215672	BB648365	BF887288	BC086482	AK142306	AK156345	AK137507	CB522550	BB865764	BY751007	BB659498	CN682136	CN682253	CF165306	AW153178	DN626023	BF270756	DU475651	BB865389	AG891620	CI643011	CI540319	CI538130	CD657296	DR005834	AL698932	DA248756	DA189529	BP315317	BP310850	BP282535	BP280749	BP353869	BP351331	DB067714	DA328439	DB134417	DB091786	DA375387	DB071966	BM146766	DB060916	DB070018	BX643165	CT346760	ID
•		BB64	BF887288 CM4-TN014	BC086482 Mus muscu	AK142306 Mus muscu	AK156345 Mus muscu		CB522550 UI-M-GH0-	BB865764 BB865764	BY751007 BY751007	BB659498 BB659498	CN682136 E0160C01-	CN682253 E0161G12-	CF165306 B0754H03-	AW153178 se36h11.y	DN626023 UI-M-EY0-	BF270756 GAEb000	DU475651 109841573	BB865389 BB865389	AG891620 Oryza sat	CI643011 CI643011	CI540319 CI540319	CI538130 CI538130	CD657296 AGENCOURT	DR005834 TC116022	. AL698932 DKFZp686F	DA248756 DA248756	DA189529 DA189529			BP282535 BP282535		BP353869 BP353869	BP351331 BP351331	DB067714 DB067714	-	DB134417 DB134417	. DB091786 DB091786	DA375387 DA375387	DB071966 DB071966	BM146766 TCAAP1E73	DB060916 DB060916	DB070018 DB070018	BX643165 DKFZp781P	CT346760 Sus scrof	Description

ALIGNMENTS

ACCESSION	DEFINITION	Locus	CT346760	RESULT 1
CT346760	Sus scrofa genomic clone CH242-41J6, genomic survey sequence.	CT346760 499 bp DNA linear GSS 03-NOV-2005		

SOURCE ORGANISM REFERENCE AUTHORS VERSION KEYWORDS RESULT 2 BX643165 밁 8 FEATURES COMMENT COMMENT DEFINITION SOURCE REFERENCE KEYWORDS VERSION ACCESSION Snoo TITLE JOURNAL AUTHORS Matches Best Local Similarity Query Match JOURNAL TITLE ORGANISM source 361 ATAATCCAGGCCATAGGAAAAAATATTTCCACTTAAGATGA 401 10 ATACTCCAGGACAAGGGAAAAACTATTTCCAATGATGATGA 50 DKFZp781P1622_r1 781 (synonym: hlcc4) Homo sapiens cDNA clone
DKFZp781P1622_5', mRNA sequence.
BX64165 CT346760.1 GI:79916147 GSS. EST. humquery@sanger.ac.uk Unpublished This sequence was generated from the SP6 end of BAC 41J6. 41J6 is part of the CHORI-242 BAC Library created by P. de Jong. Further Fobo,G., Han,M. and Wiemann,S.
EST (Koehrer,K., Beyer,A., Mewes,H.W., Weil,B., Amid,C., et al.)
Unpublished (2003) Humphray, S.J., Plumb, R.W. and Durham, J.L. Sus scrofa Koehrer, K., Beyer, A., Mewes, H.W., Weil, B., Amid, C., Osanger, A., Catarrhini; Hominidae; Homo.
1 (bases 1 to 230) Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Primates; Haplorrhini; Homo sapiens BX643165.1 GI:34477498 BX643165 details: http://www.sanger.ac.uk/Projects/S_scrofa/. Submitted (01-NOV-2005) The Sanger Institute, Wellcome Trust Genome Campus, Hinxton, Cambridgeshire, CB10 15A, UK. E-mail enquiries: Direct Submission Mammalia; Eutheria; Laurasiatheria; Cetartiodactyla; Suina; Suidae; Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Homo sapiens (human) Contact: MIPS Ingolstaedter Landstr.l, D-85764 Neuherberg, Germany (bases 1 to 499) Conservative /mol_type="genomic DNA" /db_xref="taxon:9823" /note="vector pTARBAC1.3_BamHI tissue_type="White blood cells"/ /clone="CH242-41J6" /organism⇔"Sus scrofa" Location/Qualifiers 56.4%; 0; Mismatches Score 28.2; Pred. No. 34; DB 19; 8; Indels Length 499; 0 Gaps 0;

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RESULT 3
DB070018
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AUTHORS
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DEFINITION
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Best Local Similarity
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                     JOURNAL
                                                                                                    TITLE
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     This is the 5' sequence of the clone insert Clone from S. Wiemann, Molecular Genome Analysis, German Cancer Clone from S. Wiemann, Molecular Genome Analysis, German Cancer Research Center (DKF2); bmail s. Wiemann@dkfz- heidelberg.de; sequenced by BMFZ (Biomedical Research Center at the Heinrich-Heine-University, Duesseldorf/Germanny) within the CDNA sequencing consortium of the German Genome Project. No sl sequence available. This clone (DKFZp781P1622) is available at the RZPD in Berlin.
                                                                                              Yamashita,R., Yamamoto,J., Sekine,M., Tsuritani,K., Wakaguri,H., Ishii,S., Sugiyama,T., Saito,K., Isono,Y., Irie,R., Kushida,N., Yoneyama,T., Otsuka,R., Kanda,K., Yokoi,T., Kondo,H., Wagatsuma,M., Murakawa,K., Ishida,S., Ishibashi,T. Takahashi,Fujii,A., Tanase,T., Nagai,K., Kikuchi,H., Nakai,K., Isogai,T. and Sugano,S. Diversification of Transcriptional Modulation: Large-scale
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DB070018 TESTI4 Homo sapiens cDNA clone TESTI4012418 5', mRNA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DB070018
DB070018.1 GI:83492925
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Berlin-Charlottenburg, GERMANY; Email: clone@rzpd.de.
                  Genome Res. 16 (1), 55-65 (2006)
                                                                                                                                                                                                                                                      1 (bases 1 to 531)
Kimura, K., Wakamatsu, A., Suzuki, Y., Ota, T., Nishikawa, T.,
                                                                                                                                                                                                                                                                                                            Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                               Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
                                                                                                                                                                                                                                                                                                                                                                                        Homo sapiens
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                                                                                                                                                                                                                                                                                                                                    Mammalia; Eutheria; Euarchontoglires; Primates; Haplorrhini;
16344560
                                                   Promoters of Human Genes
                                                                         [dentification and Characterization of Putative Alternative
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/note="Vector: pSportl_Sfi; Site_1: SfiIA; Site_2: SfiIB;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            /dev_stage="adult"
/lab_host="DH10B"
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Pred. No. 42;
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COMMENT

Contact: Takao Isogai

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search,
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57 E
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Title: Perfect score: Sequence:	Run on:
US-10-751-113-3 50 1 tgccatgtgatactccaggaactatttccaatgatgatga 50	March 7, 2007, 14:13:51; Search time 155 Seconds (without alignments) 603.583 Million cell updates/sec

Scoring table: IDENTITY_NUC Gapop 10.0 , Gapext 1.0

Total number of hits satisfying chosen parameters: 2807332

1403666 seqs, 935554401 residues

Searched:

Minimum DB seq length: 0
Maximum DB seq length: 200000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

O	O						Result No.
7	σ	v	4	w	N	٢	No.
24.2	24.2	26.2	26.2	26.4	27.8	27.8	Score
48.4	48.4	52.4	52.4	52.8	55.6	55.6	1
48.4 524032	48.4 524032	4860	4621	157822	6835	6760	Query Match Length DB ID
w	w	ω	w	w	w	ω	Ba
US-09-949-016-16929	US-09-949-016-16928	US-09-445-353E-1	US-09-125-635-9	US-09-949-016-16723	US-09-125-635-1	US-09-949-016-4981	Query Match Length DB ID
Sequence 16929, A	Sequence 16928, A	Sequence 1, Appli	Sequence 9, Appli	Sequence 16723, A	Sequence 1, Appli	Sequence 4981, Ap	Description

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Sequence 5, Appli Sequence 5, Appli Sequence 5, Appli Sequence 5, Appli Sequence 5, Appli Sequence 5, Appli	5, 51, 13, 51, 51, 51, 51, 51, 51, 51, 51, 51, 51	67 12 13	Sequence 15729, A Sequence 12008, A Sequence 17056, A Sequence 9, Appli Sequence 9, Appli	14346, 14347, 12086, 17390, 17391, 3, Appl	Sequence 16930, A Sequence 16931, A Sequence 14340, A Sequence 14341, A Sequence 14342, A Sequence 14343, A Sequence 14344, A Sequence 14344, A

ALIGNMENTS

US-09-949-016-4981 ; Sequence 4981, Application US/09949016 ; Patent No. 6812339

GENERAL INFORMATION:

; APPLICANT: VÉNTER, J. Craig et al.

; TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED

; TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES

THEREOF

FILE REFERENCE: CL001307
CURRENT APPLICATION NUMBER: US/09/949,016
CURRENT FILING DATE: 2000-04-14
PRIOR APPLICATION NUMBER: 60/241,755
PRIOR FILING DATE: 2000-10-20

; Sequence 16723, Application US/09949016
; Patent No. 6812339 밁 S 밁 US-09-949-016-4981 US-09-949-016-16723 US-09-125-635-1 US-09-125-635-1 SOFTWARE: FastSEQ for Windows Version 4.0 SEQ ID NO 4981 GENERAL INFORMATION: Query Match 55.6%; Score 27.8; DB 3; Length 6835; Best Local Similarity 82.1%; Pred. No. 0.93; Matches 32; Conservative 0; Mismatches 7; Indels 0 SEQ ID NO 1 Sequence 1, Application US/09125635 Patent No. 6562589 Matches Query Match Query Match 55.6%; Score 27.8; DB 3; Length 6760; Best Local Similarity 82.1%; Pred. No. 0.93; CURRENT APPLICATION NUMBER: US/09/125,635
CURRENT FILLING DATE: 1998-08-21
PRIOR APPLICATION NUMBER: 60/049,728
PRIOR FILING DATE: 1997-06-17 SOFTWARE: PatentIn Ver. 2.0 NUMBER OF SEQ ID NOS: TITLE OF INVENTION: AIB1, A novel steriod receptor co-activator FILE REFERENCE: 49944 APPLICANT: THE UNITED STATES OF AMERICA represented by THE SE NUMBER OF SEQ ID NOS: 207012 PRIOR FILING DATE: 2000-09-08 PRIOR APPLICATION NUMBER: 60/237,768 PRIOR FILING DATE: 2000-10-03 PRIOR APPLICATION NUMBER: 60/231,498 NAME/KEY: CDS LOCATION: (201)..(4463) TYPE: DNA
ORGANISM: Homo sapiens ORGANISM: Human TYPE: DNA FEATURE: LENGTH: 6835 LENGTH: 6760 Conservative 0; Mismatches 7; Indels 0 <u>,,</u> Gaps Gaps 0 0

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; NAME/KEY: CDS
; LOCATION: (110)..(4318)
US-09-125-635-9
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SOFTWARE: FastSEQ for Windows Version 4.0 SEQ ID NO 16723
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              GENERAL INFORMATION:
APPLICANT: VENTER, J. Craig et al.
TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED
TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Sequence 9, Application US/09125635 Patent No. 6562589
                                                                                                                                                                                                                                                SOFTWARE:
SEQ ID NO 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Matches
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  Best Local Similarity 79.9 Matches 31; Conservative
                                              Query Match
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PRIOR APPLICATION NUMBER: 60/231,498
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CURRENT FILING DATE: 2000-04-14
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            PRIOR APPLICATION NUMBER: 60/241,755
PRIOR FILING DATE: 2000-10-20
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                                                                                                                                                                                                                                                                                                                                 CURRENT APPLICATION NUMBER: US/09/125,635
CURRENT FILING DATE: 1998-08-21
PRIOR APPLICATION NUMBER: 60/049,728
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                                                                                                                                                                                                                                                                                                                                                                                                    FILE REFERENCE:
                                                                                                                                                                                                                                                                                                                                                                                                                         APPLICANT: THE UNITED STATES OF AMERICA represented by THE SE
TITLE OF INVENTION: AIB1, A novel steriod receptor co-activato
                                                                                                                                                                                                                                                                                            NUMBER OF SEQ ID NOS: 12
                                                                                                                                                                                                                                                                                                                 PRIOR FILING DATE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          TYPE: DNA
                                                                                                                                                      ORGANISM: Mus musculus FEATURE:
                                                                                                                                                                                                    TYPE: DNA
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52.4%; Score 26.2; Di
79.5%; Pred. No. 3.4;
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                                           DB 3; Length 4621;
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OM nucleic - nucleic search, using sw model

Run on: March 7, 2007, 14:51:58; Search time 481 Seconds (without alignments) 604.921 Million cell updates/sec

Title:

Sequence: Perfect score: US-10-751-113-3
50
1 tgccatgtgatactccagga.....actatttccaatgatgatga 50

Scoring table: IDENTITY_NUC
Gapop 10.0 , Gapext 1.0

11084925 seqs, 2909669241 residues

Total number of hits satisfying chosen parameters: 22169850

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 10% Maximum Match 100% Listing first 45 summaries

Published_Applications_NA_New: *

Database :

10: 11: 12: | EMC_Celerra_SIDSJ/ptodata/1/pubpna/US08_NEW_PUB.seq:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US06_NEW_PUB.seq:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US07_NEW_PUB.seq:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US07_NEW_PUB.seq:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US09_NEW_PUB.seq:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US09_NEW_PUB.seq:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US10_NEW_PUB.seq:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US10_NEW_PUB.seq1:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US10_NEW_PUB.seq3:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US10_NEW_PUB.seq4:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US10_NEW_PUB.seq6:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US10_NEW_PUB.seq6:*
| EMC_Celerra_SIDSJ/ptodata/1/pubpna/US11_NEW_PUB.seq6:*
|

SUMMARIES

and is derived by analysis of the total score distribution.

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed,

Result No. Score Query Match Length DB IJ

Description

Sequence 58886, A Sequence 42217, A	13 US-11-491-125A-58886 13 US-11-491-125A-42217	5298 22244	49.2	24.6	44	
	-11-266-748A-1616	789	50.4	25.2	43	n
	US-11-266-	789	50.4	25.2	42	
Sequence 63140, A	5 US-09-815-264-63140	20784	52.0	26	41	n
12582,	US-11-491-	20779	52.0	26	40	a
Sequence 9660, Ap	13 US-11-491-125A-9660	14959	52.0	26	39	O
O	13 US-11-491-125A-31020	3718	52.0	26	38	a
	7 US-10-940-774-16723	157822	52.8	26.4	37	
J	5 US-09-815-264-74332	44601	55.2	27.6	36	
Sequence 39406, A	13 US-11-491-125A-39406	42163		27.6	35	
Sequence 65963, A	5 US-09-815-264-65963	42163	•	27.6	34	
Sequence 20056, A	w	33842		27.6	33	
Sequence 59918, A	_	33842	55.2	27.6	32	
Sequence 26230, A	w	30100	55.2		31	a
Sequence 69888, A	5 US-09-815-264-69888	25882	55.2	27.6	30	
	ω	23605	•	27.6	29	
Sequence 70240, A	5 US-09-815-264-70240	23604	55.2	27.6	28	
Sequence 21915, A	13 US-11-491-125A-21915	19831	•	27.6	27	
Sequence 61400, A	5 US-09-815-264-61400	19831	55.2	27.6	26	
	c	17450			25	
Sequence 41656, A	-	17359	•		24	
	w	15173			23	a
	c	15173			22	o
2859	ω	7819	55.2	27.6	21	o
Sequence 70329, A	- :	7819	•	27.6	20	n
	•	6347	•	27.6	19	
Sequence 32347, A	-	5747		.7	18	
Sequence 29558, A	w	4056		.7	17	a
•	c	4056	55.2	7	16	o
	US-11-491-	3206			15	
	w	1742	55.2	•	14	
		1742	٠	27.6	13	
		1501	55.2	•	12	
_	US-09-815-264-16262	1501	55.2		11	
e 5955	w	1415	55.2	27.6	10	a
		1415	•	•	9	Ω
	5 US-09-815-264-40949	1113		27.6	œ	
	US-09-815-2	1052	ŗ	7	7	n
•	ω	1021	55.2	27.6	6	n
10247	US-09-8	1021	5	•	5	O
	US-11-283-329-153	7935	5	7.	4	
56675	US-11-266-	7935		7	w	
e 155,	-283	7923	55.6	27.8	ν	
Sequence 4981, Ap	8 US-10-940-774-4981	6760	55.6	27.8	_	
,		1111111				!

ALIGNMENTS

US-10-940-774-4981

Sequence 4981, Application US/10940774
Publication No. US20070037165A1
GENERAL INFORMATION:
APPLICANT: VENTER, J. Craig et al.

US-11-283-329-155 ; Sequence 155, Application US/11283329 ; Publication No. US20060134670A1 문 S US-10-940-774-4981 THEREOF SEQ ID NO 155 LENGTH: 7923 TYPE: DNA GENERAL INFORMATION: Best Local Similarity 82.1%;
Matches 32; Conservative SEQ ID NO 4981 Best Local Similarity Matches 32; Conserv Query Match CURRENT APPLICATION NUMBER: US/10/940,774
CURRENT FILING DATE: 2004-99-15
PRIOR APPLICATION NUMBER: 60/241,755 CURRENT APPLICATION NUMBER: US/11/283,329
CURRENT FILING DATE: 2005-11-18
PRIOR APPLICATION NUMBER: 60/629,811
PRIOR FILING DATE: 2004-11-19 APPLICANT: Piu, Fabrice
TITLE OF INVENTION: ENABLING TOOLS TO IDENTIFY LIGANDS FOR
TITLE OF INVENTION: HORMONE NUCLEAR RECEPTORS PRIOR FILING DATE: 2000-10-03
PRIOR APPLICATION NUMBER: 60/231,498 NUMBER OF SEQ ID NOS: 207012 SOFTWARE: FastSEQ for Windows Version 4.0 PRIOR FILING DATE: 2000-09-08 PRIOR APPLICATION NUMBER: 60/237,768 FILE REFERENCE: CL001307 SOFTWARE: FastSEQ for Windows Version 4.0 NUMBER OF SEQ ID NOS: 242 FILE REFERENCE: ACADIA.043A PRIOR FILING DATE: 2000-10-20 ORGANISM: Human OTHER INFORMATION: SRC-3 variant 2 FEATURE: NAME/KEY: CDS
LOCATION: (206)...(4468) ORGANISM: Homo sapiens TYPE: DNA LENGTH: 6760 Conservative 55.6%; Score 27.8; DB 14; Length 7923; 82.1%; Pred. No. 6.5; ative 0; Mismatches 7; Indels 0; Score 27.8; DB 8; Length 6760; Pred. No. 6.4; 0; Mismatches 7; Indels 0; Gaps Gaps 0 0,

TITLE OF INVENTION: POLYMORPHISMS IN KNOWN GENES ASSOCIATED TITLE OF INVENTION: WITH HUMAN DISEASE, METHODS OF DETECTION AND USES

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                                                                                                                                                                                       US-11-266-748A-56675
                                                                                                                                                                                                                                                       SEQ ID NO 56675
LENGTH: 7935
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Sequence 56675, Application US/11266748A Publication No. US20060134663A1
                                                                                            Matches
                                                                                                               Best Local Similarity
                                                                                                                                       Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                       PRIOR FILING DATE: 2004-11-03
PRIOR APPLICATION NUMBER: EP 04105485.9
PRIOR FILING DATE: 2004-11-03
PRIOR APPLICATION NUMBER: EP 04105484.2
PRIOR FILING DATE: 2004-11-03
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             PRIOR APPLICATION NUMBER: EP 04105483.4
PRIOR FILING DATE: 2004-11-03
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          PRIOR APPLICATION NUMBER: EP 04105479.2
PRIOR FILING DATE: 2004-11-03
PRIOR APPLICATION NUMBER: EP 04105482.6
PRIOR FILING DATE: 2004-11-03
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       CURRENT FILING DATE: 2005-11-03
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     TITLE OF INVENTION: Transcriptome Microarray Technology and TITLE OF INVENTION: Methods of Using the Same FILE REFERENCE: 55815-0102 (319189)
CURRENT APPLICATION NUMBER: US/11/266,748A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            PRIOR APPLICATION NUMBER: EP 04105507.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     APPLICANT: Johnston, Patrick APPLICANT: Mulligan, Karl
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  APPLICANT: Harkin, Paul
                                                                                                                                                                                                                                                                                                    SOFTWARE: PatentIn version 3.3
                                                                                                                                                                                                                                                                                                                               NUMBER OF SEQ ID NOS: 483996
                                                                                                                                                                                                                                                                                                                                               PRIOR APPLICATION NUMBER: US 60/700,293 PRIOR FILING DATE: 2005-07-18
                                                                                                                                                                                                                                                                                                                                                                                                 PRIOR FILING DATE: 2005-03-14
                                                                                                                                                                                                                                                                                                                                                                                                                    PRIOR APPLICATION NUMBER: US 60/662,276
                                                                                                                                                                                                          ORGANISM: Homo Sapiens
                                                                                                                                                                                                                                    TYPE: DNA
32; Conservative
                                                                                                               55.6%; Score 27.8; DB 14; Length 7935; 82.1%; Pred. No. 6.5;
                                                                                          0; Mismatches
                                                                                            7;
                                                                                            Indels
                                                                                            ..
                                                                                            Gaps
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; Sequence 153, Application US/11283329 ; Publication No. US20060134670A1 ; GENERAL INFORMATION:

US-11-283-329-153

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